

Genomic Analysis of Carbon Source Metabolism of *Shewanella oneidensis* MR-1: Predictions versus Experiments†

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Genomic sequences have been used to find the genetic foundation for carbon source metabolism in *Shewanella oneidensis* MR-1. Annotated *S. oneidensis* MR-1 gene products were examined for their sequence similarity to enzymes participating in pathways for utilization of carbon and energy as described in the BioCyc database (<http://www.biocyc.org/>) or in the primary literature. A picture emerges that relegates five- and six-carbon sugars to minor roles as carbon sources, whereas multiple pathways for utilization of up to three-carbon carbohydrates seem to be present. Capacity to utilize amino acids for carbon and energy is also present. A few contradictions emerged in which enzymes appear to be present by annotations but are not active in the cell according to physiological experiments. Annotations are based on close sequence similarity and will not reveal inactivity due to deleterious mutations or due to lack of coordination of regulation and transport. Genes for a few enzymes known by experiment to be active are not found in the genome. This may be due to extensive divergence after duplication or convergence of function in separate lines in evolution rendering activities undetectable by sequence similarity. To minimize false predictions from protein sequences, we have been conservative in predicting pathways. We did not predict any pathway when, although a partial pathway was seen it was composed largely of enzymes already accounted for in any other complete pathway. This is an example of how a biochemically oriented sequence analysis can generate questions and direct further experimental investigation.

Tens of *Shewanella* species have been isolated and named, and their phylogenetic relationships have been determined. *Shewanella* bacteria are aquatic organisms found in nature in marine ocean settings, in marshes, and in riverine and lake settings. They are gammaproteobacteria, mesophilic heterotrophic facultative anaerobes (22, 25). Although rich medium promotes faster growth, *Shewanella* species grow on minimal medium supplemented with a few nutrients, with lactate as a carbon source and any of a number of electron acceptors such as fumarate (25). *Shewanellae* are particularly distinguished by their ability to use many compounds as terminal electron acceptors in anaerobic respiration. The species whose genome we are studying is *Shewanella oneidensis*, and the strain is MR-1. It is one of a collection of bacteria isolated from sediment of Lake Oneida, N.Y. (18).

The taxonomic position of *Shewanella* bacteria has been refined over time. Early names applied to the organism were *Pseudomonas putrefaciens* and *Alteromonas putrefaciens*. *Shewanella* was recognized as a species in 1985 (15). Classification by 16S RNA sequences revealed its relationship to other aquatic organisms (25). Most recently, among *Alberomonas*-like bacteria, the genus *Shewanella* was placed within a new family, *Shewanellaceae* (10, 11).

Some of the interest in the organism stems from its ability to reduce metals and metal oxides in the environment. This has raised the possibility that shewanellae could serve as decon-

taminating agents in the environment (2, 18, 19, 24). Also, these bacteria can cause food spoilage and can act as opportunistic pathogens (3, 12) and thus are of some interest to the food industry and medicine.

Recently the *S. oneidensis* MR-1 genome was fully sequenced and its gene products were annotated (9). The early annotation has been extended since (6) and continues to be studied (13). In this work, we have placed predicted enzymes in pathways of intermediary metabolism for carbon and energy utilization in order to gain a picture from the genetic point of view of the metabolic capacities of the bacterium and to relate them to current experimental knowledge (22).

Information from the genome sequence was used to predict the presence of enzymes of carbon source metabolism. Before doing sequence comparisons, we identified fused genes in the *S. oneidensis* MR-1 genome and divided them so all gene sequences encode single proteins only (23). The list of all *S. oneidensis* MR-1 unimodular protein sequences has been compared to the protein sequences of 107 other organisms using the Darwin AllAllDb program (7). We have described previously the particular suitability of the Darwin analysis for sequence annotation (14).

Unlike the situation for most microorganisms whose genomes have been sequenced, there is a modest body of experimental information on some of the phenotypic characteristics of *S. oneidensis* MR-1. Thus, we have the opportunity to relate sequence-based predictions for *Shewanella* to existing experimental information about the organism. Do the sequence annotations for enzymes reflect experimentally known metabolic characteristics? In a few cases, experimentally derived information does not agree with the predictions. In other cases,

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information on expression would be required to relate phenotypic information to genomic results.

Some cautions and modest improvements of methods for pathway prediction have emerged in the process of genomic annotation of central metabolism of a *Shewanella* strain. Annotations of protein sequences are only predictions. We realize that there are limits to the accuracy of predicting metabolic properties by sequence comparisons. We describe steps taken to eliminate some sources of artifact from the analysis. We minimize predictions of pathways when evidence is weakened by the multiple uses of some enzymes in a cell.

Relating gene and protein sequences of any one organism to proteins of other organisms can only tell us about similarities or variations on metabolic themes already known to us through experimental work in other organisms. No completely new enzymatic functions or pathways will be revealed by current methods of gene annotation.

MATERIALS AND METHODS

Analysis of protein sequence similarities. Pairwise sequence alignments and scores were generated using the AllAllDb program of Darwin (Data Analysis and Retrieval With Indexed Nucleotide/peptide sequence package), version 2.0, developed at the ETHZ in Zurich, Switzerland (<http://cbg.inf.ethz.ch>) (7). Maximum likelihood alignments are generated with an initial global alignment by dynamic programming (Smith and Waterman algorithm) followed by dynamic local alignments (Needleman and Wunsch algorithm). A single scoring matrix is used for these steps. After the initial alignment, the scoring matrix is adjusted to fit the approximate distance between each protein pair to produce the minimum Pam value. Pam units are defined as the numbers of point mutations (base pair differences) per 100 residues. The final report includes Pam distances and variances. Darwin's ability to apply scoring matrices according to the distance between each protein pair ensures a data set of highly accurate similarity calculations (Pam scores) for distantly as well as closely related protein pairs. While the closely related homologs are mainly used for annotation purposes, the identification of distantly related proteins are valuable in finding divergent but related protein functions. The Darwin algorithms and use of multiple substitution matrices have been evaluated in relation to other sequence analysis approaches and have been given high credit for sensitivity and performance (20).

Genomic sequences for 107 microorganisms were obtained from the NCBI RefSeq web site (www.ncbi.nih.gov/RefSeq/) by ftp (<ftp://ftp.ncbi.nih.gov/genomes/Bacteria/>). Sequences of all proteins of the 107 genomes, predicted and known, were compared to sequences of all *S. oneidensis* MR-1 proteins. Data for *S. oneidensis* MR-1 genes were from the NCBI RefSeq database NC_004347.1 except that with further study some annotations have changed to greater specificity (unpublished data). A tab-delimited text table listing all genes for predicted enzymes of *S. oneidensis* MR-1 is available (see Table S1 in the supplemental material). For enzymes having sequence-similar alignments with homologs of at least 83 residues and occupying more than 45% of the sequence of both proteins, Pam values are reported. Table columns give the gene identification (ID), gene name, enzyme name or partial information, GenBank ID for the best sequence match, organism with best match, and Pam value of best match. EC numbers are included for the enzymes addressed in the paper as well as PubMed IDs for experimentally verified functions.

For the work reported here, sequence pairs were extracted from the totality of less stringent data collected: those pairs that had alignment lengths of at least 83 amino acids and distances of 125 Pam units or less. We chose the length requirement of 83 residues as it improves the significance of the sequence alignments for the more distantly related protein pairs (1). The requirement for at least 83 residues also avoids a class of commonly occurring protein domains smaller than 83 residues that appear widely in many otherwise unrelated proteins (such as small binding sites for a type of substrate, cofactor, or regulator). In addition, for this study we removed proteins directly involved in horizontal gene transfer (IS proteins, transposases, and known prophage components) from the data set.

Pathway reconstruction and vetting incomplete pathways. *S. oneidensis* MR-1 proteins annotated as enzymes were examined one by one and placed in known microbial pathways. Pathways that were experimentally determined in another organism were deemed to be present in *S. oneidensis* MR-1 by the presence of

sequence similarity for all component enzymes. Pathways for which not all enzymes were found required careful inspection. It is a fact that some enzymes occur in more than one pathway. In the cases of pathways only partially filled with homologs, we noted which of the enzymes present were already being used in other complete pathways in the cell. These have little or no significance in establishing the presence of the second pathway. Evidence for the presence of enzymes unique to the second pathway is necessary before designating them as present. We did not lower thresholds of similarity in an extended attempt to find enzymes for the holes in such tenuous pathways. Following this rule eliminated many seemingly partial pathways that we believe are not in fact present in *S. oneidensis* MR-1.

Comparison with experimental information. We have compared experimental results on central metabolism of *S. oneidensis* MR-1 with the predictions based on sequence similarity to known enzymes. Besides published work, unpublished results have kindly been made available to us as a private communication from K. E. Nealon (noted in the text as K.N.) and extensive Biolog data have been shared by J. Klappenbach and J. Tiedje. Phenotypic results, published and unpublished, are not completely consistent for this organism from different laboratories at different times, but there is agreement on the major characteristics.

RESULTS AND DISCUSSION

Sequence similarities. We identified the coding sequences (CDS) of *S. oneidensis* MR-1 that are similar in sequence to proteins in any one of the 107 genomes searched using Darwin analysis (7). A complete list of the organisms searched can be found in Table S1 in the supplemental material. Of a total of 4,325 genes coding for proteins in the *S. oneidensis* MR-1 genome, 3,382 CDS for proteins have matches by sequence similarity with Pam values up to 125 to at least one ortholog in the 107 genomes examined. The data collected are as follows: the ID of each *S. oneidensis* MR-1 protein, an abbreviation of the name of the organism, and the IDs of orthologous proteins of similar sequence, the length of all proteins, the start and end residues of both members of each alignment, the percentage of total protein aligned for each member of a pair, and the name of the orthologous protein. For this study, data for the category of enzymes were extracted.

Most closely related organisms. To identify the organisms most closely related to *S. oneidensis* MR-1 among those tested, we counted the number of proteins having matches with a Pam value less than 75 (a stringent threshold). The results, listed in column 1 of Table 1, identify *Yersinia* and *Vibrio* spp. as most similar to *S. oneidensis* MR-1. Both *Yersinia* and *Vibrio* spp. are aquatic organisms like *Shewanella*. Both are major pathogens, the meaning of which is not clear at present as shewanellae are known only to be opportunistic pathogens. In this context, the proteins of the opportunistic pathogen *Pseudomonas aeruginosa* and the pathogen *Pasteurella multocida* have many similarities to those in *S. oneidensis* MR-1. Nevertheless, it seems clear that the aqueous habitats (marine, riverine, and estuarine) of the high-ranking *Vibrio* and *Yersinia* spp. may confer the most important physiological similarities that are reflected in the level of genome sequence similarity.

As another measure of degree of relatedness, the proteins having the closest similarity to each of the *S. oneidensis* MR-1 proteins were listed by organism and counted. The result, column 2 of Table 1, shows the number of "best hits" to *S. oneidensis* MR-1 proteins. Again *Vibrio* and *Yersinia* spp. ranked higher and *Escherichia coli* species ranked lower than they did when ranked by total number of closely related homologs. Note, however, that when orthologs in closely related strains of a species are counted—in this case, *E. coli* and

TABLE 1. Similarity of genes in other genomes to those of *Shewanella oneidensis* MR-1

Organism	No. of similar genes ^a	No. of best hits ^b
<i>Yersinia pestis</i>	1,943	189
<i>Vibrio cholerae</i>	1,423	522
<i>Shigella flexneri</i>	1,342	83
<i>Escherichia coli</i>		
CFT073	1,204	42
O157:H7	1,173	43
K-12	1,135	16
<i>Erwinia carotovora</i>	1,118	103
<i>Salmonella enterica</i>		
Serovar Typhimurium	1,115	109
Serovar Typhi	1,104	16
<i>Pseudomonas aeruginosa</i>	1,087	192
<i>Pseudomonas syringae</i>	976	101
<i>Chromobacterium violaceum</i>	776	62
<i>Pasteurella multocida</i>	764	45

^a Number of hits per genome with Pam values below 75.^b Number of proteins for which the lowest Pam value is in this organism.

Salmonella enterica serovars—one must be aware that proteins closely related in all strains will be counted only once as the best match to a *Shewanella* protein, yet all are highly significant. Thus, low scores by close relatives are not indicative of genetic distance as the marginally lower similarities are not counted in the ranking. Nevertheless, *Vibrio* and *Yersinia* species far outstrip the other organisms tested.

Although *E. coli* strains ranked low in numbers of best matches to *S. oneidensis* MR-1 coding sequences, nevertheless the *E. coli* data were vital to interpretation of the results. *E. coli* among all organisms tested had the most information on gene products that have been experimentally characterized, not simply predicted by sequence similarity.

Metabolism pathways—overview. The subset of orthologs with the best Pam scores from any of the genomes as well as all significant orthologs in *E. coli* K-12, whether the best or not, were collected for all the enzymes of central metabolism. We grouped enzymes by pathway, some enzymes appearing more than once if present in more than one pathway. We found, based on sequence similarity, that *S. oneidensis* MR-1 like *E. coli* has complete pathways for the biosynthesis of all amino acids, nucleotides, and cofactors. However, utilization of compounds as carbon and energy sources was more restricted than in *E. coli*, and in some instances the pathways were not those of *E. coli* but of other organisms such as *P. aeruginosa*.

Carbon source and energy metabolism by respiration. Homologous genes for enzymes of main carbon utilization and central metabolism are shown in Fig. 1. Convincing support for the major pathways of carbon utilization and multipurpose enzymes of central metabolism is present. Sequences are present for all enzymes of the pentose pathway, the pyruvate

dehydrogenase complex, all enzymes of the tricarboxylic acid cycle, the glyoxylate bypass, the Entner-Doudoroff pathway, major anaplerotic reactions, and other enzymes of central carbohydrate metabolism. Note that by sequence similarity an essential enzyme of glycolysis, 6-phosphofructokinase, is not found. In agreement, it is known experimentally that *S. oneidensis* MR-1 does not use glucose as a carbon and energy source (Biolog; K.N.). The critical glycolytic enzyme 6-phosphofructokinase was not found in cell extracts (22). Consistent with the inability to use glucose for growth, we find that the sequences for enzymes for the feeder reactions that convert six-carbon carbohydrates to glucose that are present in *E. coli* are not found in *S. oneidensis* MR-1 (see Table S2 in the supplemental material). Among *E. coli* enzymes used in the conversion of 15 compounds, we only detected *S. oneidensis* MR-1 homologs for galactokinase and ribokinase. The presence of these two kinases does not reflect utilization of galactose or ribose for carbon and energy; rather they are conversions in the course of intermediary metabolism. The predicted enzyme composition of *S. oneidensis* MR-1 in this respect—shy on enzymes for utilization of five- and six-carbon carbohydrates—is consistent with known phenotypic characterization (25).

The existence of multiple copies of genes or genes for isozymes might reflect a gene dosage effect supporting heavy use of these particular enzymes. We note that isozymes are not present in *S. oneidensis* MR-1 for five- and six-carbon metabolism as they are in *E. coli*, but do exist for glyceraldehyde-3-phosphate dehydrogenase, an enzyme that is vital to central metabolism at the three-carbon level (see Table S3 in the supplemental material). Also in *S. oneidensis* MR-1, different from *E. coli*, large numbers of phosphotransferase system (PTS) genes for transport of simple sugars are not present. Only one set of PTS genes is found in *S. oneidensis* MR-1, whereas *E. coli* has 41 genes encoding PTS proteins for a variety of four-, five-, and six-carbon carbohydrates.

Instead of six-carbon sugars, *S. oneidensis* MR-1 prefers three-carbon carbohydrates for growth. Experimentally, L-lactate, pyruvate, and acetate are among compounds utilized as sources of carbon and energy (22, 25; Biolog data). Agreeing with the experiment, genes are present for the enzymes for metabolism of pyruvate and acetate and other two- and three-carbon molecules (Fig. 2). However, the genetic picture seemingly does not agree for L-lactate, which does in fact support growth, but no homolog for a currently sequenced bacterial L-lactate dehydrogenase is found in *S. oneidensis* MR-1. A homolog for an unrelated fermentative D-lactate dehydrogenase gene is present, although *S. oneidensis* MR-1 is not considered a fermentative organism and is unable to ferment glucose (22, 25). Since *S. oneidensis* MR-1 is usually grown with L-lactate as a carbon and energy source, the absence of a homolog for L-lactate dehydrogenase is puzzling. Perhaps there is a different kind of lactate dehydrogenase not represented in current sequence databases. We note that two genes, designated by locus tags SO1520 and SO1521, are located adjacent to the lactate permease gene. One codes for an iron-sulfur protein, and the product of the other contains a domain for binding flavin adenine dinucleotide and has similarity to a glycolate dehydrogenase. Could these CDS represent a different kind of prokaryotic L-lactate dehydrogenase, a dimer that

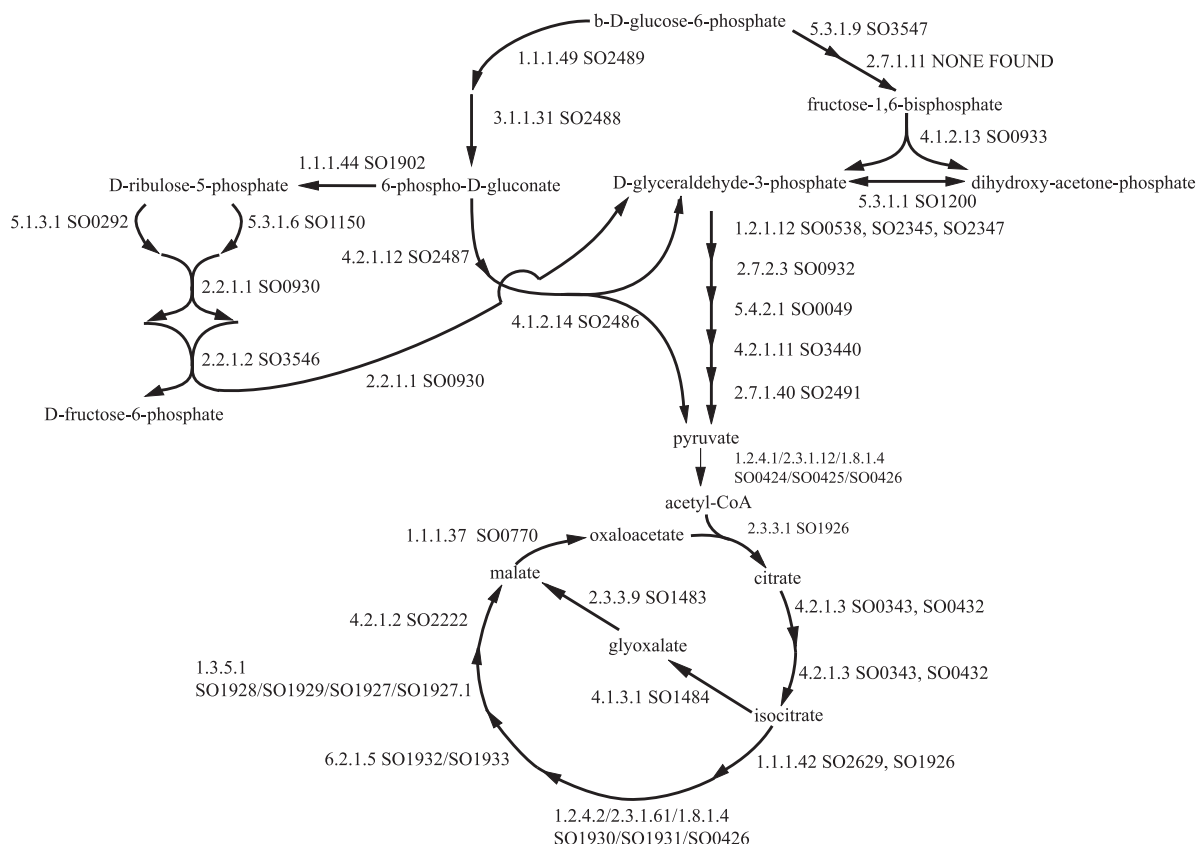


FIG. 1. Carbon source metabolism in *S. oneidensis* MR-1. The intermediary metabolic steps of glycolysis, tricarboxylic acid (TCA) cycle, glyoxalate bypass, the pentose phosphate, and the Entner-Doudoroff pathways are shown. Enzymes are shown by their EC number followed by the *S. oneidensis* locus tag (SO number) of the gene predicted to encode the respective activity. The function predictions were based on sequence similarity to proteins with experimentally verified functions. Predicted isozymes are shown as SO numbers separated by a comma. Enzyme complexes are indicated by SO numbers separated by a forward slash. Predictions were made for all steps, except for that of 6-phosphofructokinase, EC 2.7.1.11.

contains iron sulfur center(s) and flavin adenine dinucleotide bound to separate subunits? Experimental exploration of the possibility could be fruitful.

Although reports of ability to grow on propionate are mixed (22; K.N.; Biolog), on the basis of sequence similarity, support is found. Propionate could be metabolized by a variation on the 2-methyl citrate pathway in which 2-methyl citrate is converted to 2-methyl-*cis*-aconitate by the AcnD-PrpF combination, a complex 2-methyl citrate dehydratase. In this respect, *S. oneidensis* MR-1 is like *P. aeruginosa* and *Vibrio cholerae*, not like *E. coli*, which has the PrpD version of the enzyme (8).

S. oneidensis MR-1 uses a C-1 compound, formate, as an energy source (22). Standing out from a plethora of genes concerning formate are genes for three formate dehydrogenase operons and one hydrogenase operon found in the genome. The three clusters are similar to those known for three-subunit formate dehydrogenase enzymes that participate in respiration, using either oxygen as an electron acceptor or nitrate. The three clusters are SO0101 to -0103 plus the accessory protein product of SO0107; SO4509 to -4511, and SO4513 to -4515 plus the accessory protein product of SO4503. The hydrogenase SO2097, -2098, and -2099 genes code for the subunits of a quinone-reactive type of hydrogenase known to work together with respiratory formate dehydrogenases. Adja-

cent to the hydrogenase operon is a contiguous set of genes coding for the hydrogenase accessory proteins (SO2089 to SO2096). Absence of the fermentative type of formate dehydrogenase in complex with hydrogenase to make the formate hydrogen lyase complex is consistent with the observation that *S. oneidensis* MR-1 is not a fermentative organism. The genes encoding for pyruvate formate-lyase and its activator were found (SO2912 and -2913). Although this enzyme usually is associated with fermentative metabolism, it may be supplying *S. oneidensis* with C-1 (formate) units.

The CO₂ produced by formate oxidation cannot be fixed by *S. oneidensis* MR-1 as no homologs are found for any known enzymes for CO₂ fixation. Consideration has been given to the possibility *S. oneidensis* MR-1 is a facultative methylotroph with the cyclic C-1 serine pathway. Some of the enzymes for the cyclic serine pathway are present but, by stringent sequence comparisons, not all. An alternative to the classical serine pathway that uses only enzymes of general metabolism has been proposed (22). However, this version may not be able to effect net incorporation of C-1 moieties into metabolism. Since all of the enzymes in this scheme are also required for other pathways, their presence in MR-1 is ambiguous.

As to the classical C-1 cyclic serine pathway, almost all needed genes are present in *S. oneidensis* MR-1. It appears to

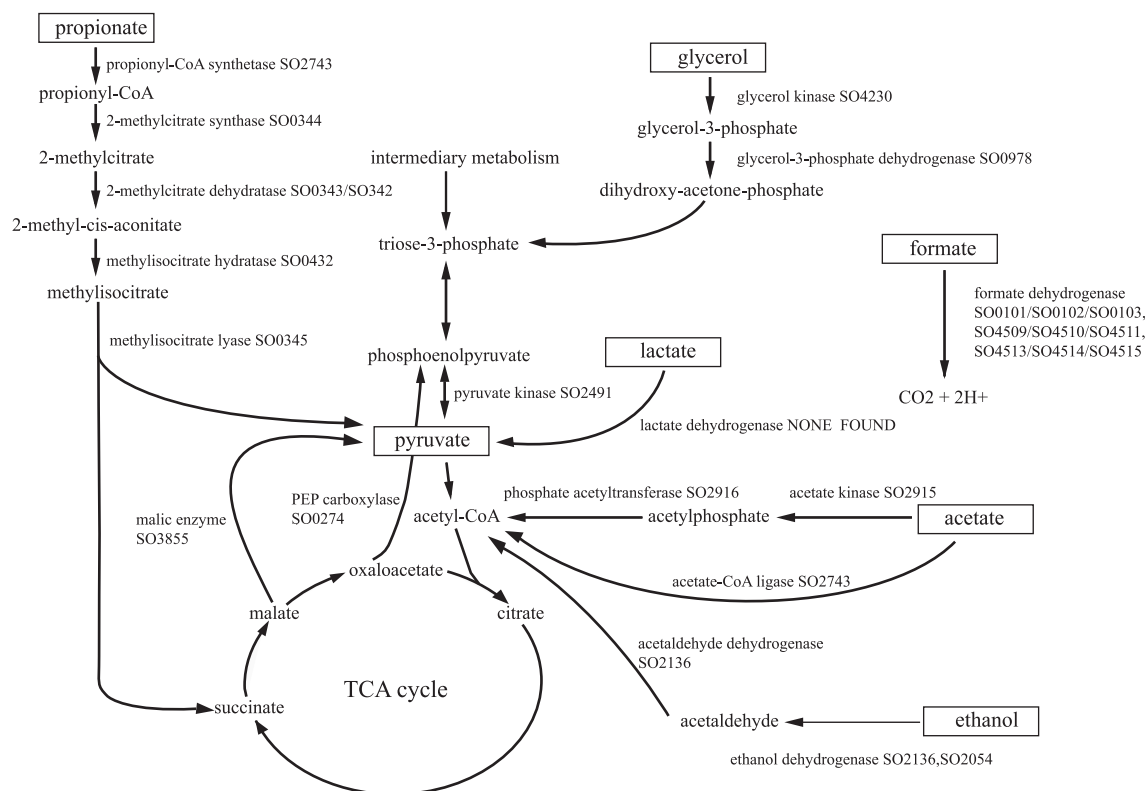


FIG. 2. Enzymes in *S. oneidensis* MR-1 for utilization of one-, two-, and three-carbon compounds. Enzymes involved in the degradation of compounds with one carbon (formate), two carbons (acetate, ethanol), and three carbons (propionate, glycerol, lactate, and pyruvate) are shown by their names and predicted *S. oneidensis* locus tags (SO number). The function predictions were based on sequence similarity to proteins with experimentally verified functions. Predicted isozymes are shown as SO numbers separated by a forward slash. Enzyme complexes are indicated by SO numbers separated by a forward slash. TCA, tricarboxylic acid.

be able to activate both formate and formaldehyde substrates by enzymes of tetrahydrofolate metabolism, converting them into the derivative *N*-5,10-methylenetetrahydrofolate (methyleneTHF) (4, 16). The methyleneTHF enters the cyclic C-1 serine pathway by donating the C-1 moiety to combine with glycine producing serine, resulting ultimately in capture of carbon as acetyl-coenzyme A (CoA) and regeneration of the C-1 acceptor molecule glycine for the next cycle of capture. There are two unique enzymes that could identify the pathway: malate-CoA ligase and malyl-CoA lyase (Fig. 3). Searching for sequences for the two unique enzymes in this pathway did not reveal homologs in *S. oneidensis* MR-1 within the threshold of significance applied in this work. However, we note that enzymes are present for similar reactions such that broad substrate specificity could enable the enzymes to encompass the activities of the two missing reactions. The succinyl-CoA ligase (SO1932 and SO1933) might serve as the malyl-CoA lyase if it also had the ability to bind malate. As a possible substitute for the malyl-CoA lyase, the *S. oneidensis* MR-1 genome has a close homolog of the gene for 3-hydroxy-3-methylglutaryl-CoA lyase (SO1893). Could this enzyme, if it has broad substrate specificity, also serve as the malyl-CoA lyase? 3-Hydroxy-3-methylglutarate and malate are chemically related compounds. Experimental characterization of the proteins of SO1893 and SO1932 to -1933 could establish whether either possibility is the case. Otherwise *Shewanella* spp. could employ an as yet

unknown pathway as a recycling mode for capturing carbon when growing on formate or formaldehyde as a carbon source (5). Biochemical investigation is needed.

Given its evident preference for C-3 and C-1 carbon sources, *S. oneidensis* MR-1 will depend on anaplerotic reactions, the glyoxylate pathway, gluconeogenesis, and the nonoxidative pentose pathway reactions to generate the intermediates required for biosynthetic metabolism. Not mentioned above are the critical anaplerotic enzymes, all of which seem to be present: phosphoenolpyruvate carboxykinase, phosphoenolpyruvate carboxylase, and malic enzyme.

Ability to utilize amino acids as carbon sources is also widespread in *S. oneidensis* MR-1 (21). Sequence similarity was found to the genes coding for the enzymes of the pathways for utilization of amino acids as they are known in *E. coli*. In addition, in *S. oneidensis* MR-1, some of the amino acids are broken down by pathways found in other organisms. These are shown in Fig. 4. The ability to utilize histidine is not by the pathway found in *E. coli* K-12 but is like the transformations in *Bacillus subtilis*. The ability to utilize aromatic amino acids is not by the pathway in *E. coli*, but has similarities to part of the protocatechuate pathway as found in pseudomonads. Genes for the enzymes of the latter steps of the protocatechuate meta-pathway are present in *S. oneidensis* MR-1, but only in the steps after the intermediate γ -carboxymuconolactone. Utilization of aromatic compounds may have unique early steps in

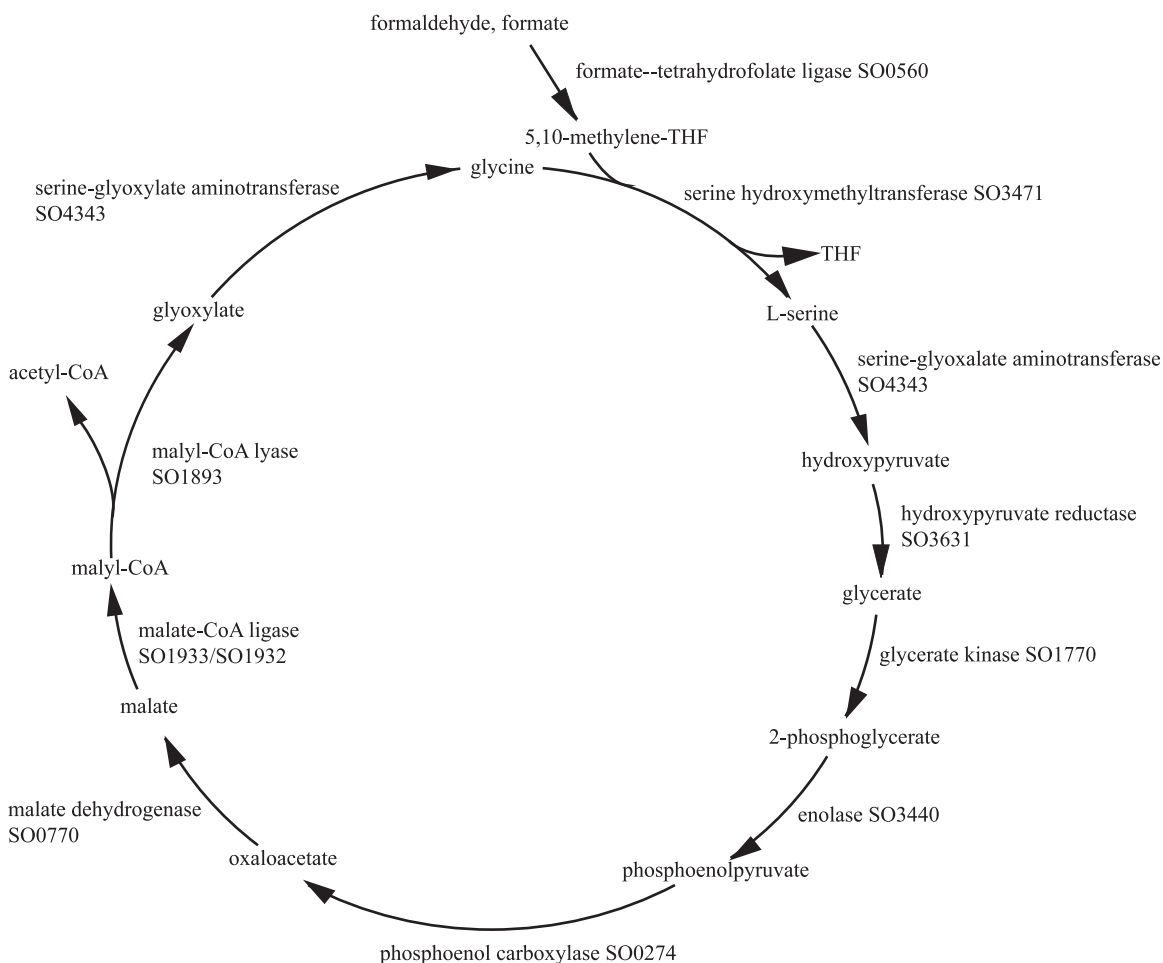


FIG. 3. Classical C-1 serine pathway enzymes in *S. oneidensis* MR-1. Prediction of *S. oneidensis* MR-1 enzymes involved in the growth and assimilation of one-carbon compounds via the serine pathway are shown. Function predictions were made based on sequence matches to proteins with experimentally verified functions. Predicted enzyme complexes are indicated by SO numbers separated by a forward slash. Significant sequence matches to the two key enzymes in the pathway, malyl-CoA lyase (EC 4.1.3.24) and malate-CoA ligase (EC 6.2.1.9), were not found. Tentative assignments for these reactions were made to *S. oneidensis* MR-1 gene products catalyzing similar reactions.

S. oneidensis MR-1 that join the known pathway at the carboxymuconolactone step. *S. oneidensis* MR-1 also degrades leucine, isoleucine, and valine not simply by transamination as in *E. coli* but rather by pathways used in pseudomonads. Methionine is converted to 2-oxobutanoate as in pseudomonads.

Even though enzymes for utilization of many amino acids seem to be present by sequence analysis, as measured by Biolog data, single amino acids do not serve as carbon sources for *S. oneidensis* MR-1. However, these Biolog results show that many dipeptides composed of many combinations of pairs of amino acids do register positively. The easier permeability and lower specificity of dipeptide transport systems compared with single-amino-acid transporters might account for the difference in results. Homologs for several dipeptide-specific ABC transport proteins are found in the genome, and there are numerous homologs of dipeptidases that could hydrolyze the dipeptides taken up and make available the monomeric amino acids for utilization.

Fatty acids are also used as carbon and energy sources. Homologs for genes coding for enzymes for breakdown of fatty

acids are found. There are multiple copies of some of the key enzymes in these pathways. Other compounds found by Biolog testing to be utilized as carbon sources are α -ketobutyric acid, methylpyruvate, and D-lactate methyl ester, each readily convertible to mainline metabolites. Also giving positive Biolog results is *N*-acetylglucosamine, for which all sequences for the enzymes of the degradative pathway are present in the genome.

Experimental information on nucleoside utilization is interesting. Biolog and laboratory (K.N.) results for utilization of the nucleosides uridine, adenosine, 2'-deoxyadenosine, inosine, and, to a lesser extent, thymidine are positive. In contrast, the Biolog results for the bases cytosine, thymine, adenine, and guanine are poor to negative. We see that sequences for nucleoside hydrolases are present in *S. oneidensis* MR-1 that could split the nucleosides to their bases and ribose-5-phosphate for further utilization. However, since Biolog data indicate that the bases are not utilized, evidently it is primarily the ribose-5-phosphate moiety that is supporting growth when

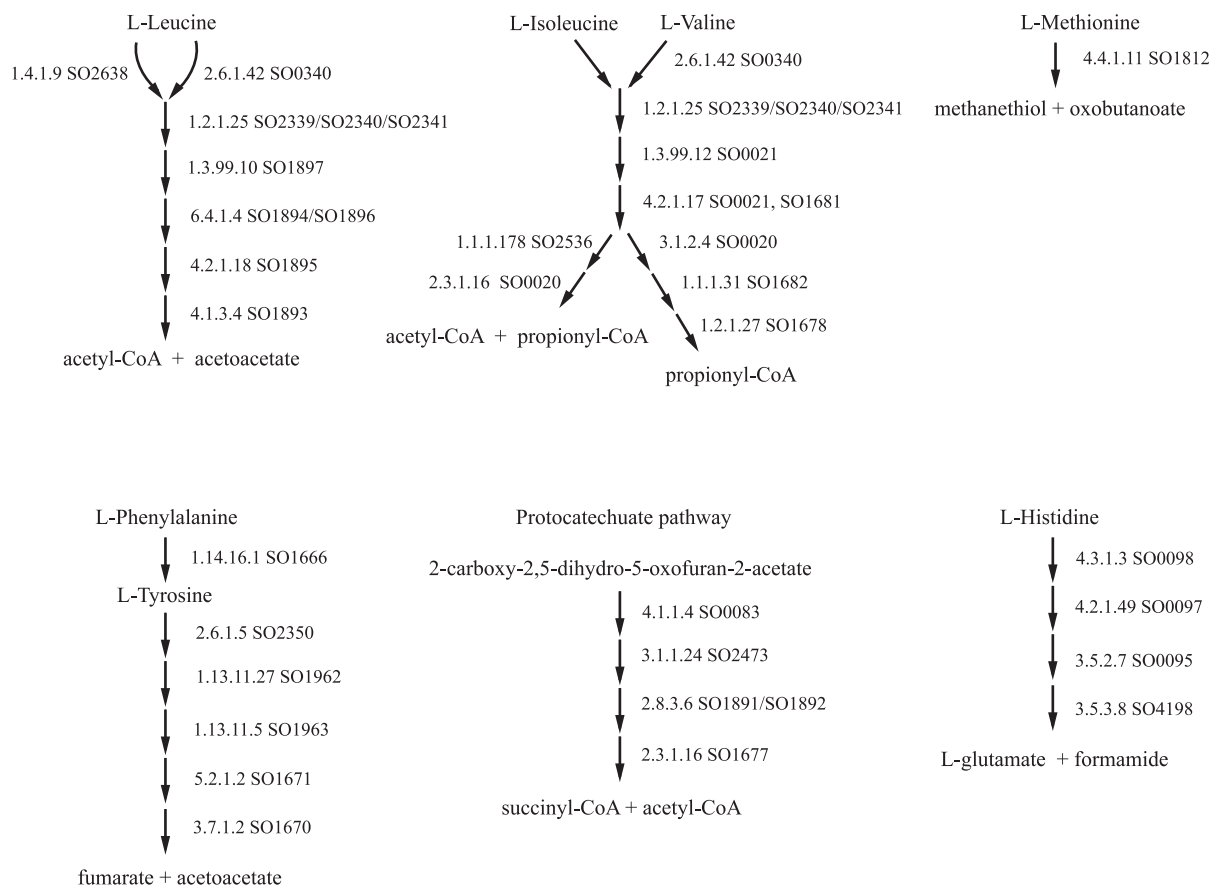


FIG. 4. Amino acid utilization in *S. oneidensis* MR-1 by other than *E. coli* pathways. *S. oneidensis* MR-1 enzymes predicted to be involved in the degradation of amino acids are shown by their locus tags (SO numbers) and the EC number of the reaction. Gene predictions are based on sequence similarities to known enzymes. Predicted isozymes are shown as SO numbers separated by a comma. Enzyme complexes are indicated by SO numbers separated by a forward slash. Only pathways not utilized by *E. coli* are included in the figure.

nucleosides are provided. Again, further experimentation is required to establish this interpretation.

Aerobic and anaerobic respiration. With respect to respiratory electron transfer pathways, electron transfer elements seem to be emphasized in the *S. oneidensis* MR-1 genome. Pathways for synthesis of heme, siroheme, menaquinone, and ubiquinone are intact. *S. oneidensis* MR-1 is believed to contain 42 cytochrome *c*-type proteins (17), whereas by comparison there are only 5 in *E. coli*.

Shewanella spp. are well known for their use in anaerobic respiration of many electron acceptors other than oxygen, the most unique being metals and metallic compounds (2, 18, 25). In agreement, genes have been found by sequence similarity to code for reduction of nitrate, nitrite, sulfate, sulfur, trimethylamine *N*-oxide (TMAO), dimethyl sulfoxide, and fumarate reductase enzymes corresponding to many of the metabolites used as anaerobic respiration acceptors. Enzymes for reduction of Fe ions are known, but reductases for the numerous metallic compounds that serve as receptors have yet to be identified. However, there is no dearth of possibilities in the genome sequences. We found many candidate uncharacterized oxidoreductases.

Fermentation. *S. oneidensis* MR-1 does not ferment (22, 25), yet many enzymes of mixed acid fermentation seem to be

present in the genome (Fig. 5). In fermentation, choices include multiple paths from lactate or pyruvate to ethanol or acetate and conversion of acetate to acetyl-CoA or acetylphosphate. Other pathways lead from formate to production of H₂ and CO₂. Although all enzymes of mixed acid fermentation are present, there is no evidence it does exist. Support for a pathway does not exist if all enzymes are known to be active in other pathways (Table 2). In some cases, such as with the cyclic C-1 serine pathway, there is at least one unique enzyme in the pathway. Designation of the pathway then hangs on the presence or absence of any unique member. Table 2 gives examples of pathways that use many or all of its enzymes that are present in other pathways. Note that all of the enzymes used in fermentation pathways are also used in other known pathways. Thus, there is no indication from genomic analysis whether fermentation is used or not by this organism.

Biosynthesis. The emphasis in this survey is on utilization of sources of carbon and energy, yet in the course of the work, the presence of homologs in the genome of enzymes of biosynthesis of basic building blocks was noted. Although *S. oneidensis* MR-1 is frequently grown with supplements to increase the growth rate, the organism is able to grow on an unsupplemented minimal medium. In agreement, based on sequence similarity, *S. oneidensis* MR-1 has complete biosynthetic capacity

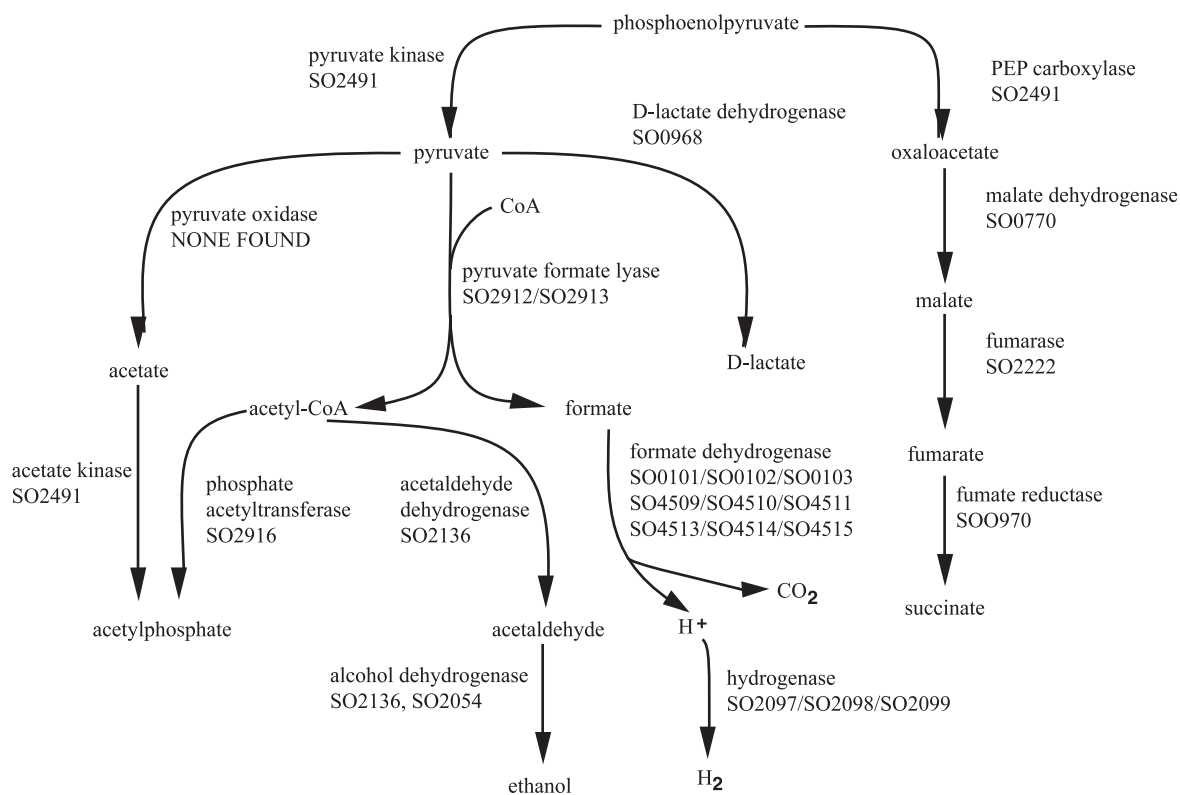


FIG. 5. Enzymes of mixed acid fermentation in *S. oneidensis* MR-1. Sequence matches for *S. oneidensis* MR-1 gene products (SO numbers) to the enzymes used by *E. coli* for mixed acid fermentation are shown. Significant similarities to all enzymes except pyruvate oxidase (EC 1.2.2.2) were found. Predicted isozymes are separated by commas. Reactions predicted to be carried out by enzyme complexes are shown by SO numbers separated by forward slashes.

for basic building blocks and cofactors. Sequences are found for enzymes similar to those in *E. coli* for biosynthesis of all amino acids, purines, pyrimidines, and growth factors. The essential precursor metabolites that are needed for biosynthetic activities of the cell are as follows: glucose-6-P, fructose-6-P, ribose-5-P, erythrose-4-P, triose phosphate, 3-phosphoglycerate, phosphoenolpyruvate, pyruvate, acetyl-CoA, 2-oxoglutarate, succinyl-CoA,

oxalacetate, and sedoheptulose-7-phosphate. In *S. oneidensis* MR-1, all enzymes required to generate these precursors are supported by genomic sequences.

Summary of general considerations of pathway prediction. We have emphasized that there are caveats for pathway prediction which we applied in this work. A common approach to connecting information about enzyme homologs with metabolic pathways is to list the enzymes that seem to be present for any and all pathways. By assessing pathways one by one, if all enzymes are present one asserts the pathway is present in the organism. If not all enzymes are present, there is a temptation to look for the missing enzymes, relaxing thresholds of sequence similarity in an effort to complete the pathway list. However, it is important first to ask whether any of the enzymes in a partial pathway are used in other pathways in the cell. If all enzymes of a pathway with “holes” are used in other pathways, then there is no evidence for existence of the partial pathway. For some pathways, there is at least one unique enzyme in the pathway that is not used in any other known pathway in the cell. If a homolog for the unique enzyme is found, then it does seem safe to assert that pathway is present, even if not all other enzymes have yet been found. Illustrations are given in Table 2. In the classical cyclic C-1 serine pathway, two enzymes are unique. However, for mixed acid fermentation, all enzymes are also used in general metabolism; none are unique to the fermentation. Therefore, one cannot know without experimentation whether any of the fermentations occur.

TABLE 2. Some pathways that use enzymes in common with other pathways

Pathway 1	No. of enzymes		Pathway 2
	Common	Unique	
C-1 serine pathway	2		Tricarboxylic acid cycle
	1		Glycolysis/gluconeogenesis
	1		Glyoxylate bypass
	1		Central metabolism
		2	
Mixed acid fermentation	3		Tricarboxylic acid cycle
	3		Glycolysis
	4		Central metabolism
		0	
Valine utilization	2		Leucine utilization
		5	
Glycolysis	8		Gluconeogenesis
		1	

In asserting a metabolic capability of any kind by genomic analysis, one needs to keep in mind that not all pathways in the biological world are yet known, so failure to find standard ways to convert one substance to another need not mean the biological capability is absent in the organism. An example is evidence in *S. oneidensis* MR-1 for only part of the protocatechuate pathway of aromatic compound degradation.

In all predictions of enzyme activity, it goes without saying that sequence similarity does not guarantee biological activity. Mutations may be present that produce an inactive enzyme product. We have tried to demur with assertions like “appears to be present” since we cannot know whether a homolog that is not identical to an active enzyme is in fact capable of activity.

Concluding remarks. Identification of homologs to enzymes of metabolism has allowed us to assemble a picture of central carbon metabolism in *S. oneidensis* MR-1. Utilization of carbohydrates larger than three carbons is not well supported by sequence similarity to appropriate genes and enzymes. Utilization of three-carbon carbohydrates and smaller has genetic support. Capability for utilization of amino acids and fatty acids as carbon sources is also present. Basic biosynthetic enzymes are present. Most pathways were similar to those in *E. coli* even though the closest homologs by sequence analysis were not often found in *E. coli*. But these pathways presumably are highly similar in the most closely related gammaproteobacteria, *Vibrio* spp. and *Yersinia* spp. Exceptions were found for some pathways for utilization of amino acids that were similar to those of pseudomonads or bacilli.

There are opportunities for connecting study of the physiology and metabolism of *S. oneidensis* MR-1 in the laboratory with questions of the kind raised here about operation of particular pathways and the properties of particular enzymes. Each of the questions posed here provides an opportunity for investigation that has the potential of yielding new knowledge about microbial metabolism. This would seem to be a useful outcome of sequence similarity analysis of genomic DNA.

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